

# eam and Deep UV Exposure of PMMA Based Resists -Identical or Different Chemical Behavior?

A. Uhl \*b, J. Bendig \*, J. Leistner \*, U. Jagdhold b, J. Bauer b

Humboldt University of Berlin, Institute of Chemistry, Hessische Str. 1-2, 10115 Berlin, Germany

Institute for Semiconductor Physics, Walter-Korsing-Str. 2, 15230 Frankfurt (Oder), Germany

### Abstract

The chemical reactions and the dissolution properties of homopolymeric PMMA and a P(MMA-co-MAA) copolymer were investigated during DUV (KrF, 248 nm) and e-beam exposure. The chain scission reaction was analyzed using GPC. The polymer degradation reaction is very similar at both exposure procedures. In both cases a bimodal and, later, a multimodal character of the molecular weight distribution is observed.

### Introduction

produce images with excellent profiles. chain scission efficiency. Although, resists of the PMMA type require very high exposure doses, but, they resists show a low sensitivity as a consequence of the low absorptivity (e.g. at  $\lambda_{
m sc}=248$  nm) and low (P(MMA-co-MAA)), characterized by an approximately four times higher sensitivity, have been used as positive resists in electron beam tithography for a long time. At deep UV exposure the PMMA based Poly(methyl methacrylate) (PMMA) and its derivative Poly(methyl methacrylate -co- methacrylic acid)

electron beam and deep UV light using spectroscopic and chromatographic methods. In both cases, the pathway of the polymer fragmentation is globally described by a Norrish Type I cleavage forming low been done to describe the chemical changes in PMMA and its derivative caused by irradiation with possibility to predict the lithographic behavior of the resist. Therefore, on one side investigations have The description of their lithographic performance has not only been done in experimental ways but calculations have been used, too. Only the use of correct parameters in these calculations provides the show a more complex mechanism. molecular products. Details of the structure of products formed (bimodal molecular weight distribution)

and the solubility rate on the basis of our experimental results. Proceeding from the independent In this paper we check the applicability of the formula for the e-beam induced molecular weight change values for PMMA and P(MMA-co-MAA). density we consider their cometation with each other and compare experimental data with calculated experimental determination of both the solubility rate and the molecular weight on the deposited energy

the photochemistry at e-beam und DUV exposure and discuss different behavior. Compared with analogous experimental results for the deep UV degradation we show both similarities in

Homopolymeric PMMA (Polyscience) and the photoresist ARP 610 (Aliresist, Germany), dissolved in 2-methyl-glycol-acetate and 2-methyl-glycol, respectively, were spin-coated onto 4' silicon waters to a thickness of about 1 µm and baked at 170°C for 1 hour.

### E-beam exposure

The resist layers were electron beam exposed with an accelerating voltage of 20 kV over the whole wafer and different doses were applied (exposure tool ZBA 23, Jenoptik, Germany).

1452 I SPIE Val. 3333

0277-786X/98/\$10.00

The resist layers were DUV exposed using a KrF excimer laser EMG 104 MSC (Lambda Physik, Germany) and a DUV stepper NSR-2205EX12B (Nikon). The pulse energy used was 0.6 ... 3.9 mJ/cm² (EMG 104 MSC) and 0.23 mJ/cm² (DUV stepper). Different doses were applied controlled by the amount of pulses.

# UV spectroscopy

spectrometer U-3410 (Hitachi). The UV irradiation spectra of the exposed resist layers were measured on quartz substrates using a

## GPC measurements

2 ml tetrahydrofuran resp. methanol and analyzed by gel permeation chromatography (UV detection at 224 nm) using PL gel 10 mm mlx B and PL gel 100 Å columns (Knauer), PMMA reference standards (Polyscience) and tetrahydrofuran as eluent. Thereafter molecular-weight distribution curves and the weight-average molecular weight  $\overline{\mathsf{M}}_{\mathsf{w}}$  (formula 1) of each sample were determined using PSS Win GPC To perform a GPC measurement the resist was completely removed from the wafers and dissolved scientific V2.71 software.

$$\overline{M}_{W} = \frac{\sum_{i} r_{i} M_{i}}{\sum_{i} r_{i} M_{i}} \tag{1}$$

# Dissolution characteristics

dissolution at a temperature of T = wet development process, density was determined in a depending on the deposite dose presented in Tab.I. thickness as a function of the measuring ketone : Isopropanol (MIBK/I developer containing meth The dissolution rate of ARP rate was deter ਛੋਂ remainin 610

MIBIVIPA at 22 °C	Table I. Dissolution rate of ARP 610 in					gose presented in Tab.i.	thickness as a function of the exposure	the asuming and remaining resist	1 1 1	at a temperature of 1 = 64 or 110	Recone : Isopropation (wild or 17)	developer containing methyr-isobutyr	wet development process, using a	depending on the deposited energy density was determined in a standard	THE CHARGE IN THE CONTRACT OF THE CHARGE IN
F 472	3.879	3.448	3.017	2.586	2.155	1.724	1.293	0.882	0.690	0.517	0.345	0.172		Deposited energy density E [eV/nm <sup>2</sup> ]	
20 2000	10.2940	7.7270	5.3600	3.0680	2.3200	1.2586	1.0690	0.5000	0.3750	0.2820	0.1930	0.1064		Dissolution rate [nm/s]	

MIBK/IPA at 22 °C

Fig. 1 show the irradiation spectra of ARP 610 and PMMA at DUV exposure. The spectral changes at ebeam exposure are very similar.

weight distribution is shifted to lower molecular weights by increasing dose in all cases. A new character The DUV exposure dose was 0 mJ/cm $^2$   $\le$  D  $\le$  36449 mJ/cm $^2$  (ARP 610, Fig. 2b) and 0 mJ/cm $^2$   $\le$  D  $\le$  200149 mJ/cm $^2$  (PMMA, Fig. 3b). Figs. 2 and 3 show that the maximum of the molecular dose was 0 μC/cm² < D < 150 μC/cm² (ARP 610, Fig. 2a), 0 μC/cm² < D < 300 μC/cm² (PMMA, Fig. 3a) ARP 610 at e-beam and DUV exposure (1. 248 nm), respectively. The investigated range of e-beam GPC was used to investigate the dose dependence of the molecular weight distribution of PMMA and

۷

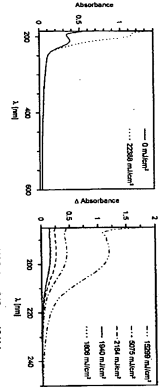


Fig. 1 Irradiation spectra during DUV exposure (laser: EMG 104 MSC;  $\lambda_{\rm ex}$  = 248 nm; 10 Hz) a; ARP 610 on quartz; b; PMMA on quartz; Δ Absorbance = Absorbance (tω) - Absorbance (t=0)

of the molecular weight distribution curve is observed with increasing dose starting with a value of D = 30µC/cm² (ARP 610, Fig. 2a and PMMA, Fig. 3a), D = 7687 mJ/cm² (ARP 630, Fig. 2b) and D = 12000 mJ/cm² (PMMA, Fig. 3b). A low molecular weight shoulder comes up with a maximum of about 2000 p/ mol. The molecular weight distribution becomes bimodal. In the range of higher doses > 80 µC/cm² and > 12000 mJ/cm², respectively, we do not observe a further shift of the higher weight maximum at M = 10000 g/mol, but its decrease to the credit of the lower weight fraction M = 2000 g/mol

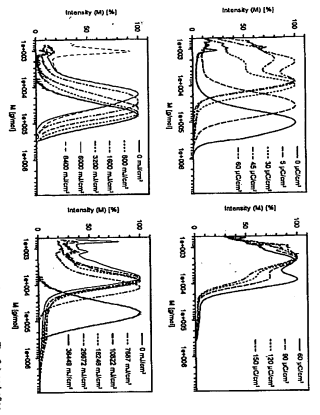


Fig. 2. GPC analytical data of ARP-810 after electron beam exposure (upper curves, Fig. 2a) and after DUV exposure (lower curves, Fig. 2b)

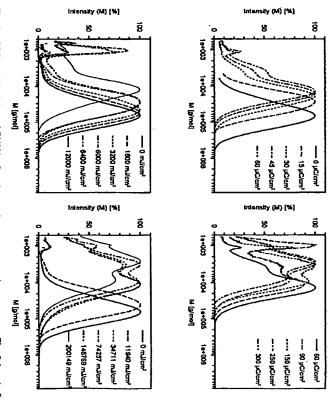


Fig. 3. GPC analytical data of PMMA after electron beam exposure (upper curves, Fig. 3a) and after DUV exposure (lower curves, Fig. 3b)

In addition the distribution becomes multimodal, and a corresponding increase of molecular weight fraction at M = 1000 g/mol and less is detected. In the case of the DUV exposed resist materials the formation of the weight fraction M = 1000 g/mol is stronger than for e-beam treated materials. The bimodal and the multimodal distribution are the result of specific chain scission. Comparing the molecular weight of ARP 610 and PMMA, the chain scission efficiency is higher for ARP 610 than for

molecular weight of one monomer unit M. = 98.4 g/mol [-(C5H6O2)0,74-(C4H6O2)0,76-corresponding to 74 % molecular weight: 28 g/mol. The following resist data were applied: density of ARP 610 1.04 g/cm deposited energy density was determined in a standard wet development process. The conversion between the dose D [µC/cm²] and the deposited energy density E [eV/nm²] was made by calculations methyl methacrylate units and 26 % methacrylic acid units in ARP 610]. The results are presented in Tab.I. using Monte Carlo methods. The used substrate data are: density of the silicon layer. behavior we measured the dissolution rate. Therefore, the dissolution rate of ARP 610 depending on the To study the influence of chain scission of the resist during electron beam exposure on the lithographic 2.33 g/cm

### 4. Discussion

chain scission of PMMA to check its applicability on the investigated resists ARP 610 and PMMA. at electron beam exposure calculations are performed using the GREENEICH model (formula 2)\* of statistic For a more comprehensive understanding of the dependence of molecular weight distribution on the dose

Ø

 $M_{\rm sc}$  fragmented weight-average molecular weight;  $M_{\rm sc}$ : weight-average molecular weight at D = 0  $\mu C/cm^2$ , g: scission efficiency; E: deposited energy density;

p: resist density (p(ARP 610) = 1.04 g/cm<sup>2</sup>; p(PMMA) = 1.1 g/cm<sup>3</sup>); N<sub>a</sub>: Avogadro's number

Applying the Greeneich model the following different scission efficiencies were calculated:
g (ARP 610) = 1.5°10° events/eV and g (PMMA) = 5°10° events/eV. The scission coefficient for ARP 610 corresponds to the value used by Greeneich. The g value of ARP 610 is about four times higher than that of PMMA, the influence of methacrylic acid units in ARP 610 leads to an increase of the scission efficiency

densities cause increasing differences between experimental and calculated MJ/M<sub>40</sub> data. For both Fig. 4. Calculated M<sub>2</sub>/M<sub>20</sub> data are in agreement with experimental data at low deposited energy densities. At higher E values (ARP 610 > 4 eV/nm², PMMA > 8 eV/nm²) increasing deposited energy resists values for the weight-average molecular weight are found to be higher by experiment than Experimentally determined Na/Nao were compared to calculated Na/Nao data. The results are shown in

The experimentally determined data of the DUV exposure are shown in Fig. 4, too.

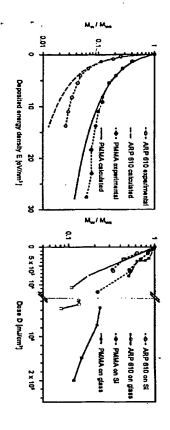
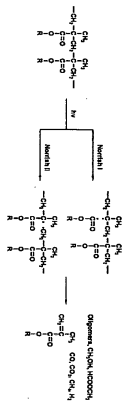


Fig. 4. Dependence of motecular weight distribution on exposure dose - left: e-beam; right: DUV

cleavage. Beside the degradated polymer the volatile products shown in the scheme are detected using GC-MS (m/z = 41, 42, 43, 44, 45, 58, 59, 60, 61, 74). During both exposure procedures within the first decomposition. The reason for the enhanced formation of these oligomers (= decamers) is still unknown. formation of the molecular weight fraction of about 1000 g/mol is favored during the second step of The mechanism of the e-beam and DUV induced chain scission is mainly characterized by a Norrish I The degradation pathways during e-beam and DUV exposure are very similar. period the molecular weight is shifted to lower values (Figs. 2 and 3) caused by the chain cleavage. The



Photochemical reactions in poly(methacrylic acid) derivative based polymers

The analytical data confirm the very similar chemical behavior of PMMA and P(MMA-co-MAA) at e-beam and DUV exposure. The chemical pathway is characterized by the chain scission (positive resist) and, more and more, by the favored formation of specific fractions (bimodal and multimodal distribution).

# Acknowledgment

Semiconductor Physics, for cooperation at the DUV stepper exposure experiments The authors thank the Fonds der Chemischen Industrie for financial support and Mr. U. Haak, Institute for

### 7. References

- M. Hatzakis, J. Vac. Sci. Technol. 16 (1979) 1984; W. Moreau, M. Merrit, W. Moyer, M. Hatzakis, J. Vac. K. U. Pohl, F. Rodriguez, Y. M. N. Namaste, S. K. Obendorf, ACS Symp. Ser. 266 (1984) 323; A. Uhl, J. Bendig, J. Leistner, U. Jagdhold, L. Bauch, M. Bottcher, J. Vac. Sci. Technol., in preparation.
- Sci. Technol. 16 (1979) 1989.
- 3 K. Jain, Excimer Laser Lithography, SPIE, Bellingham 1990, p. 139-145
- J. S. Greeneich and T. van Duzer, IEEE Trans. Electron Devices, ED 21 (1974) 286